SCORE Search Results Details for Application 10716359 and Search Result 20061227... Page 1 of 2

<!--StartFragment-->

GenCore version 5.1.9

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OM protein - protein search, using sw model

December 27, 2006, 21:00:21; Search time 199 Seconds Run on:

(without alignments)

1346.376 Million cell updates/sec

US-10-716-359-16 Title:

Perfect score: 3104

1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2589679 segs, 457216429 residues

2589679 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 '

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 500 summaries

Database : A Geneseq 8:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:* 6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

9: geneseqp2005s:*

10: geneseqp2006s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| | | · O | | | | |
|--------|-------|-------|--------|----|----------|--------------------|
| Result | | Query | | | | |
| No. | Score | Match | Length | DB | ID | Description |
| 1 | 3104 | 100.0 | 586 | 2 | AAY05956 | Aay05956 Human cel |
| 2 | 3104 | 100.0 | 586 | 5 | ABG95140 | Abg95140 Human onc |
| 3 | 3100 | 99.9 | 586 | 5 | ABG95132 | Abg95132 Human onc |
| 4 | 3087 | 99.5 | 586 | 3 | AAB11357 | Aab11357 Human p63 |
| 5 | 3087 | 99.5 | 586 | 5 | ABB74989 | Abb74989 Human p53 |
| 6 | 3087 | 99.5 | 586 | 5 | ABP61909 | Abp61909 Human lun |
| 7 | 3087 | 99.5 | 586 | 7 | ADH36992 | Adh36992 Human lun |
| ٠ 8 | 3087 | 99.5 | 586 | 8 | ADM56795 | Adm56795 Human lun |
| 9 | 3087 | 99.5 | 586 | 8 | ADN04517 | Adn04517 Antipsori |
| 10 | 3087 | 99.5 | 586 | 9 | ADU98487 | Adu98487 Human p53 |
| 11 | 3087 | 99.5 | 586 | 9 | AEB10295 | Aeb10295 Cancer re |
| 12 | 3087 | 99.5 | 586 | 9 | AEE06149 | Aee06149 p53 tumor |
| 13 | 3080 | 99.2 | 586 | 2 | AAY41032 | Aay41032 Human lun |
| 14 | 3080 | 99.2 | 586 | 3 | AAB11317 | Aab11317 Human lun |
| 15 | 3080 | 99.2 | 586 | 5 | ABB74949 | Abb74949 Human lun |
| | | | | | | |

| | | • | | | | | |
|----------|------|------|------------|---|----------|------------|-----------|
| 16 | 3080 | 99.2 | 586 | 5 | ABP61869 | Abp61869 | Human lun |
| 17 | 3080 | 99.2 | 586 | 7 | ADE53447 | Ade53447 | Human lun |
| 18 | 3080 | 99.2 | 586 | 7 | ADH36806 | Adh36806 | Human lun |
| 19 | 3080 | 99.2 | 586 | 8 | ADM56609 | Adm56609 | Human lun |
| 20 | 3080 | 99.2 | 586 | 8 | ADN89653 | Adn89653 | Human lun |
| 21 | 3080 | 99.2 | 586 | 9 | ADU98301 | Adu98301 | Lung tumo |
| 22 | 3080 | 99.2 | 586 | 9 | AEB10109 | Aeb10109 | Cancer re |
| 23 | 3080 | 99.2 | 586 | 9 | AEE05963 | Aee05963 | Human lun |
| 24 | 3078 | 99.2 | 586 | 7 | ADA28428 | Ada28428 | p53 tumou |
| 25 | 3077 | 99.1 | 586 | 7 | ADA28242 | Ada28242 | Human lun |
| 26 | 3066 | 98.8 | 586 | 2 | AAY05962 | Aay05962 | Mouse cel |
| 27 | 3033 | 97.7 | 641 | 2 | AAY45247 | | Human p51 |
| 28 | 3033 | 97.7 | 641 | 2 | AAY05953 | | Human cel |
| 29 | 3033 | 97.7 | 641 | 4 | AAB82129 | | Human pro |
| 30 | 3033 | 97.7 | 641 | 5 | ABG95142 | | Human onc |
| 31 | 3033 | 97.7 | 687 | 7 | ADL22695 | | Human dis |
| 32 | 3029 | 97.6 | 641 | 3 | AAB11358 | | Human p63 |
| 33 | 3029 | 97.6 | 641 | 5 | ABG95136 | | Human onc |
| 34 | 3029 | 97.6 | 641 | 5 | ABG95128 | | Human onc |
| 35 | 3029 | 97.6 | 641 | 5 | ABB74990 | | Human p53 |
| 36 | 3029 | 97.6 | 641 | 5 | ABP61910 | | Human lun |
| 37 | 3029 | 97.6 | 641 | 7 | ADA28429 | | p53 tumou |
| 38 | 3029 | 97.6 | 641 | 7 | ADH36993 | | Human lun |
| 39 | 3029 | 97.6 | 641 | 8 | ADM56796 | | Human lun |
| 40 | 3029 | 97.6 | 641 | 9 | ADU98488 | | Human p53 |
| 41 | 3029 | 97.6 | 641 | 9 | AEB10296 | | Cancer re |
| 42 | 3029 | 97.6 | 641 | 9 | AEE06150 | | p53 tumor |
| 43 | 3025 | 97.5 | 680 | 3 | AAY50997 | | Human KET |
| 44 | 3025 | 97.5 | 680 | 3 | AAB11361 | | Human p63 |
| 45 | 3025 | 97.5 | 680 | 5 | ABB74993 | | Human p53 |
| 46 | 3025 | 97.5 | 680 | 5 | ABP61913 | | Human lun |
| 47 | 3025 | 97.5 | 680 | 7 | ADA28432 | | p53 tumou |
| 48 | 3025 | 97.5 | 680 | 7 | ADH36996 | | Human lun |
| 49 | 3025 | 97.5 | 680 | 8 | ADM56799 | | Human lun |
| 50 | 3025 | 97.5 | 680 | 8 | ABM80526 | | Tumour-as |
| 51 | 3025 | 97.5 | 680 | 8 | ADU06284 | | Novel bro |
| 52 | 3025 | 97.5 | 680 | 9 | ADU98491 | | Human p53 |
| 53 | 3025 | 97.5 | 680 | 9 | AEB10299 | | Cancer re |
| 53 54 | 3025 | 97.5 | 680 | 9 | AEE06153 | | p53 tumor |
| 55 | 2995 | 96.5 | 680 | 2 | AAY05958 | | Human cel |
| | | | | 3 | | | Rat KET p |
| 56 | 2991 | 96.4 | 680 461 | | AAY50998 | | Human p63 |
| 57 | 2421 | 78.0 | 461 | 3 | AAB11362 | | Human onc |
| 58 | 2421 | 78.0 | 461 | 5 | ABG95131 | | |
| 59 | 2421 | 78.0 | 461 | 5 | ABG95138 | | Human onc |
| 60 | 2421 | 78.0 | 461 | 5 | ABB74994 | Abb / 4994 | Human p53 |

<!--EndFragment-->

SCORE Search Results Details for Application 10716359 and Search Result 20061227... Page 1 of 2

<!--StartFragment-->

GenCore version 5.1.9

Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on: December 27, 2006, 21:01:02; Search time 306 Seconds

(without alignments)

1771.435 Million cell updates/sec

Title: US-10-716-359-16

Perfect score: 3104

Sequence: 1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 500 summaries

Database: UniProt 7.2:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| | _ | | 8 | | | | |
|----|-------|--------|-------|--------|----|--------------|--------------------|
| Re | esult | | Query | | | | |
| | No. | Score | Match | Length | DB | ID | Description · |
| - | 1 | 3033 | 97.7 | 680 | 1 | P73L HUMAN | Q9h3d4 homo sapien |
| | 2 | 3026 | 97.5 | 582 | 2 | Q3UVĪ3 MOUSE | Q3uvi3 mus musculu |
| | 3 | 3017 | 97.2 | 582 | 2 | Q5CZX0 MOUSE | Q5czx0 mus musculu |
| | 4 | 2995 | 96.5 | 680 | 1 | P73L MOUSE | 088898 mus musculu |
| | 5 | 2991 | 96.4 | 680 | 1 | P73L RAT | Q9jjp6 rattus norv |
| | 6 | 2962 | 95.4 | 582 | 2 | Q9DEC7_CHICK | Q9dec7 gallus gall |
| | 7 | 2416 | 77.8 | 565 | 2 | Q4S122 TETNG | Q4s122 tetraodon n |
| | 8 | 2344 | 75.5 | 576 | 2 | Q8JFE3 BRARE | Q8jfe3 brachydanio |
| | 9 | 2330 | 75.1 | 588 | 2 | Q8JHZ6_BRARE | Q8jhz6 brachydanio |
| | 10 | 1877 | 60.5 | 393 | 2 | Q569E5 MOUSE | Q569e5 mus musculu |
| | 11 | 1826 | 58.8 | 457 | 2 | Q8JHZ5 BRARE | Q8jhz5 brachydanio |
| | 12 | 1777 | 57.2 | 641 | 2 | Q9W664 BARBU | Q9w664 barbus barb |
| | 13 | 1774 | 57.2 | 640 | 2 | Q801Z7_BRARE | Q801z7 brachydanio |
| | 14 | 1771 | 57.1 | 640 | 2 | Q6UNX2 BRARE | Q6unx2 brachydanio |
| | 15 | 1748.5 | 56.3 | 590 | 2 | Q9JJP1_MOUSE | Q9jjp1 mus musculu |
| | 16 | 1744.5 | 56.2 | 631 | 2 | Q9JJP2_MOUSE | Q9jjp2 mus musculu |
| | 17 | 1741 | 56.1 | 636 | 1 | P73_HUMAN | 015350 homo sapien |
| | 18 | 1730.5 | 55.8 | 365 | 2 | Q98SW0_XENLA | Q98sw0 xenopus lae |
| | 19 | 1720.5 | 55.4 | 637 | 1 | P73_CERAE | Q9xsk8 cercopithec |
| | 20 | 1579 | 50.9 | 434 | 2 | Q6DG24_BRARE | Q6dg24 brachydanio |
| | 21 | 1562 | 50.3 | 514 | 2 | Q9CU77_MOUSE | Q9cu77 mus musculu |
| | 22 | 1507.5 | 48.6 | 494 | 2 | Q3UT91_MOUSE | Q3ut91 mus musculu |
| | 23 | 1481 | 47.7 | 662 | 2 | Q4S837 TETNG | Q4s837 tetraodon n |

| 24 | 1464 | 47.2 | 555 | 2 | Q5KQU6_BRARE | Q5kqu6 | brachydanio |
|----|--------|------|-----|---|--------------|-------------|-------------|
| 25 | 1459.5 | 47.0 | 497 | 2 | Q9WUJ0_MOUSE | Q9wuj0 : | mus musculu |
| 26 | 950.5 | 30.6 | 621 | 2 | Q9NGC7_MYAAR | Q9ngc7 | mya arenari |
| 27 | 912 | 29.4 | 591 | 2 | Q8T7V3_SPISO | | spisula sol |
| 28 | 912 | 29.4 | 596 | 2 | Q6WG19_SPISO | | spisula sol |
| 29 | 912 | 29.4 | 596 | 2 | Q6WG20_SPISO | Q6wg20 | spisula sol |
| 30 | 896.5 | 28.9 | 564 | 2 | Q27937_LOLFO | Q27937 | loligo forb |

SCORE Search Results Details for Application 10716359 and Search Result 20061227... Page 1 of 2

<!--StartFragment-->

GenCore version 5.1.9

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OM protein - protein search, using sw model

Run on: December 27, 2006, 21:04:46; Search time 41 Seconds

(without alignments)

1375.196 Million cell updates/sec

Title: US-10-716-359-16

Perfect score: 3104

Sequence: 1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 500 summaries

Database : PIR_80:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| | | 6 | | | | |
|--------|-------|-------|--------|----|--------|--------------------|
| Result | | Query | | | | |
| No. | Score | Match | Length | DB | ID | Description |
| 1 | 1774 | 57.2 | 640 | 2 | JC7989 | p73 protein - zebr |
| 2 | 839.5 | 27.0 | 396 | 1 | JH0631 | cellular tumor ant |
| 3 | 762 | 24.5 | 363 | 1 | A29376 | cellular tumor ant |
| 4 | 758.5 | 24.4 | 367 | 1 | S02193 | cellular tumor ant |
| 5 | 744 | 24.0 | 386 | 1 | S51648 | cellular tumor ant |
| 6 | 723 | 23.3 | 391 | 1 | S02192 | cellular tumor ant |
| 7 | 712 | 22.9 | 396 | 1 | JH0633 | cellular tumor ant |
| 8 | 707.5 | 22.8 | 393 | 1 | DNHU53 | cellular tumor ant |
| 9 | 703 | 22.6 | 393 | 1 | S06594 | cellular tumor ant |
| 10 | 702 | 22.6 | 390 | 1 | DNMS53 | cellular tumor ant |
| 11 | 700 | 22.6 | 391 | 2 | JC6193 | tumor suppressor p |
| 12 | 697.5 | 22.5 | 393 | 2 | JC6176 | tumor suppressor p |
| 13 | 688.5 | 22.2 | 381 | 2 | S38824 | cellular tumor ant |
| 14 | 259.5 | 8.4 | 77 | 2 | 146226 | cellular tumor ant |
| 15 | 152.5 | 4.9 | 925 | 2 | T19361 | hypothetical prote |
| 16 | 134.5 | 4.3 | 901 | 2 | JC6093 | dead ringer nuclea |
| 17 | 133 | 4.3 | 1081 | 2 | S66736 | transcription acti |
| 18 | 133 | 4.3 | 1520 | 1 | TVFFA | protein-tyrosine k |
| 19 | 132.5 | 4.3 | 2578 | 2 | A56922 | transcription fact |
| 20 | 130.5 | 4.2 | 2529 | 2 | A56923 | transcription fact |
| 21 | 129 | 4.2 | 963 | 2 | T40290 | hypothetical prote |
| | | | | | | |

| hypothetical prote | T41547 | 2 | 964 | 4.2 | 129 | 22 |
|--------------------|--------|---|------|-----|-------|----|
| KIAA1304 protein [| G59436 | 2 | 1051 | 4.2 | 129 | 23 |
| hypothetical prote | T15264 | 2 | 1621 | 4.2 | 129 | 24 |
| hypothetical prote | S19150 | 2 | 628 | 4.1 | 128.5 | 25 |
| hypothetical 69K p | JQ0110 | 2 | 628 | 4.1 | 126.5 | 26 |
| cell proliferation | B48666 | 2 | 2897 | 4.1 | 126 | 27 |
| cell proliferation | A48666 | 2 | 3256 | 4.1 | 126 | 28 |
| hypothetical prote | S01955 | 2 | 628 | 4.0 | 125.5 | 29 |
| hypothetical prote | T47149 | 2 | 724 | 4.0 | 125 | 30 |
| | | | | | | |

<!--StartFragment-->

GenCore version 5.1.9

Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on: December 27, 2006, 21:10:02; Search time 50 Seconds

(without alignments)

1025.859 Million cell updates/sec

Title: US-10-716-359-16

Perfect score: 3104

Sequence: 1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 500 summaries

Database: Issued Patents AA:*

1: /EMC Celerra SIDS3/ptodata/2/iaa/5_COMB.pep:*

2: /EMC Celerra SIDS3/ptodata/2/iaa/6 COMB.pep:*

3: /EMC_Celerra_SIDS3/ptodata/2/iaa/7_COMB.pep:*

4: /EMC_Celerra_SIDS3/ptodata/2/iaa/H_COMB.pep:*

5: /EMC Celerra SIDS3/ptodata/2/iaa/PCTUS_COMB.pep:*

6: /EMC Celerra SIDS3/ptodata/2/iaa/RE COMB.pep:*

7: /EMC Celerra SIDS3/ptodata/2/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | % Query Match | Length | DB | ID | Description |
|---------------|--------|---------------------|--------|----|--------------------------------|-------------------|
| 1 | 3104 | 100.0 | 586 | 2 | US-09-538-106-16 | Sequence 16, Appl |
| 2 | 3104 | 100.0 | 586 | 3 | US-09-529-583-16 | Sequence 16, Appl |
| 3 | 3087 | 99.5 | 586 | 2 | US-09-643-597-338 [.] | Sequence 338, App |
| 4 | 3087 | 99.5 | 586 | 2 | US-09-542-615A-338 | Sequence 338, App |
| 5 | 3087 | 99.5 | 586 | 2 | US-09 - 606-421B-338 | Sequence 338, App |
| 6 | 3087 | 99.5 | 586 | 2 | US-09-630-940B-338 | Sequence 338, App |
| 7 | 3087 | 99.5 | 586 | 2 | US-10-007-700-338 | Sequence 338, App |
| 8 | 3080 | 99.2 | 586 | 2 | US-09-643-597-152 | Sequence 152, App |
| 9 | 3080 | 99.2 | 586 | 2 | US-09-480-884A-152 | Sequence 152, App |
| 10 | 3080 | 99.2 | 586 | 2 | US-09-542-615A-152 | Sequence 152, App |
| 11 | 3080 | 99.2 | 586 | 2 | US-09-606-421B-152 | Sequence 152, App |
| 12 | . 3080 | 99.2 | 586 | 2 | US-09-221-107-152 | Sequence 152, App |
| 13 | 3080 | 99.2 | 586 | 2 | US-09-466-396A-152 | Sequence 152, App |
| 14 | 3080 | 99.2 | 586 | 2 | US-09-476-496A-152 | Sequence 152, App |
| 15 | 3080 | 99.2 | 586 | 2 | US-09-630-940B-152 | Sequence 152, App |
| 16 | 3080 | 99.2 | 586 | 2 | US-09-285-479-152 | Sequence 152, App |
| 17 | 3080 | 99.2 | 586 | 2 | US-10-007-700-152 | Sequence 152, App |
| 18 | 3066 | 98.8 | 586 | 2 | US-09-538-106-22 | Sequence 22, Appl |

```
586
                              US-09-529-583-22
                                                           Sequence 22, Appl
      3066
             98.8
                           3
      3033 .
             97.7
                      641
                              US-09-538-106-13
                                                           Sequence 13, Appl
20
                           2
             97.7
                      641
                              US-09-529-583-13
                                                           Sequence 13, Appl
21
      3033
                           3
                                                           Sequence 339, App
                      641
                              US-09-643-597-339
22
      3029
             97.6
                           2
                                                           Sequence 339, App
23
      3029
             97.6
                      641
                              US-09-542-615A-339
                      641
                              US-09-606-421B-339
                                                           Sequence 339, App
24
      3029
             97.6
                           2
                                                           Sequence 339, App
      3029
             97.6
                      641
                           2
                              US-09-630-940B-339
25
      3029
             97.6
                      641
                          2
                              US-10-007-700-339
                                                           Sequence 339, App
26
                                                           Sequence 342, App
      3025
             97.5
                      680
                           2
                              US-09-643-597-342
27
                                                           Sequence 342, App
      3025
             97.5
                      680
                              US-09-542-615A-342
28
29
      3025
             97.5
                      680
                              US-09-606-421B-342
                                                           Sequence 342, App
             97.5
                      680
                                                           Sequence 342, App
30
      3025
                              US-09-630-940B-342
      3025
             97.5
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                              US-10-007-700-342
                                                           Sequence 342, App
31
      2995
             96.5
                      680
                           2
                              US-09-538-106-19
                                                           Sequence 19, Appl
32
                                                           Sequence 19, Appl
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                           3
                              US-09-529-583-19
33
                                                           Sequence 343, App
34
      2421
             78.0
                      461
                           2
                              US-09-643-597-343
                                                           Sequence 343, App
35
      2421
             78.0
                      461
                              US-09-542-615A-343
                                                           Sequence 343, App
             78.0
                      461
36
      2421
                              US-09-606-421B-343
37
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             78.0
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                           2
                              US-09-630-940B-343
                                                           Sequence 343, App
                                                           Sequence 17, Appl
      2421
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                      461
                           2
                              US-09-538-106-17
38
                                                           Sequence 343, App
      2421
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                      461
                              US-10-007-700-343
39
                                                           Sequence 17, Appl
40
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             78.0
                      461
                              US-09-529-583-17
      2398
             77.3
                      461
                              US-09-538-106-23
                                                           Sequence 23, Appl
41
                                                           Sequence 23, Appl
      2398
             77.3
                      461
                              US-09-529-583-23
42
             75.7
                      516
                              US-09-643-597-344
                                                           Sequence 344, App
      2350
43
             75.7
                                                           Sequence 344, App
      2350
                      516
                              US-09-542-615A-344
44
                                                           Sequence 344, App
45
      2350
             75.7
                      516
                           2
                              US-09-606-421B-344
      2350
             75.7
                      516
                           2
                              US-09-630-940B-344
                                                           Sequence 344, App
46
                      516
                                                           Sequence 14, Appl
47
      2350
             75.7
                           2
                              US-09-538-106-14
      2350
             75.7
                      516
                          2
                              US-10-007-700-344
                                                           Sequence 344, App
48
             75.7
                          3
49
      2350
                      516
                              US-09-529-583-14
                                                         . Sequence 14, Appl
50
      2327
             75.0
                      555
                          2
                              US-09-538-106-20
                                                           Sequence 20, Appl
```

<!--EndFragment-->

OM protein - protein search, using sw model

December 27, 2006, 21:21:46; Search time 186 Seconds Run on:

(without alignments)

1459.376 Million cell updates/sec

US-10-716-359-16 Title:

Perfect score: 3104

1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

2097797 segs, 463214858 residues Searched:

Total number of hits satisfying chosen parameters: 2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100%

용

Listing first 500 summaries

Database : Published Applications AA Main:*

1: /EMC Celerra SIDS3/ptodata/2/pubpaa/US07 PUBCOMB.pep:*

2: /EMC Celerra SIDS3/ptodata/2/pubpaa/US08 PUBCOMB.pep:*

3: /EMC Celerra SIDS3/ptodata/2/pubpaa/US09 PUBCOMB.pep:*

4: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US10A_PUBCOMB.pep:*

5: /EMC Celerra SIDS3/ptodata/2/pubpaa/US10B_PUBCOMB.pep:*

6: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US11_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | Query Match | Length | DB | ID | Description |
|---------------|---------------|----------------|--------|----|--------------------|-------------------|
| 1 | 3104 | 100.0 | 586 | 4 | US-10-716-359-16 | Sequence 16, Appl |
| 2 | 3104 | 100.0 | 586 | 5 | US-10-469-469-306 | Sequence 306, App |
| 3 . | 3104 | 100.0 | 586 | 6 | US-11-109-831-16 | Sequence 16, Appl |
| 4 | 3100 | 99.9 | 586 | 5 | US-10-469-469-290 | Sequence 290, App |
| 5 | 3087 | 99.5 | 586 | 3 | US-09-735-705-338 | Sequence 338, App |
| , 6 | 3087 | 99.5 | 586 | 3 | US-09-850-716A-338 | Sequence 338, App |
| . 7 | 3087 | 99.5 | 586 | 3 | US-09-897-778-338 | Sequence 338, App |
| 8 | 3087 | 99.5 | 586 | 4 | US-10-007-700-338 | Sequence 338, App |
| 9 | 308· 7 | 99.5 | 586 | 4 | US-10-117-982-338 | Sequence 338, App |
| 10 | 3087 | 99.5 | 586 | 4 | US-10-313-986-338 | Sequence 338, App |
| 11 | 3087 | 99.5 | 586 | 5 | US-10-775-972-338 | Sequence 338, App |
| 12 | 3087 | 99.5 | 586 | 5 | US-10-922-124-338 | Sequence 338, App |
| 13 | 3087 | 99.5 | 586 | 5 | US-10-623-155-338 | Sequence 338, App |
| 14 | 3080 | 99.2 | 586 | 3 | US-09-735-705-152 | Sequence 152, App |
| İ5 | 3080 | 99.2 | 586 | 3 | US-09-850-716A-152 | Sequence 152, App |
| 16 | 3080 | 99.2 | 586 | 3 | US-09-897-778-152 | Sequence 152, App |
| 17 | 3080 | 99.2 | 586 | 3 | US-09-466-396A-152 | Sequence 152, App |
| 18 | 3080 | 99.2 | 586 | 4 | US-10-007-700-152 | Sequence 152, App |
| 19 | 3080 | 99.2 | 586 | 4 | US-10-117-982-152 | Sequence 152, App |

| 20 | 3080 | 99.2 | 586 | 4 | US-10-313-986-152 | Sequence | 152, App |
|----|------|------|-----|---|--------------------|----------|----------|
| 21 | 3080 | 99.2 | 586 | 5 | US-10-775-972-152 | Sequence | 152, App |
| 22 | 3080 | 99.2 | 586 | 5 | US-10-922-124-152 | Sequence | 152, App |
| 23 | 3080 | 99.2 | 586 | 5 | US-10-623-155-152 | Sequence | 152, App |
| 24 | 3066 | 98.8 | 586 | 4 | US-10-716-359-22 | Sequence | 22, Appl |
| 25 | 3066 | 98.8 | 586 | 6 | US-11-109-831-22 | Sequence | 22, Appl |
| 26 | 3033 | 97.7 | 641 | 4 | US-10-716-359-13 | Sequence | 13, Appl |
| 27 | 3033 | 97.7 | 641 | 5 | US-10-469-469-310 | Sequence | 310, App |
| 28 | 3033 | 97.7 | 641 | 6 | US-11-109-831-13 | Sequence | 13, Appl |
| 29 | 3029 | 97.6 | 641 | 3 | US-09-735-705-339 | Sequence | 339, App |
| 30 | 3029 | 97.6 | 641 | 3 | US-09-850-716A-339 | Sequence | 339, App |
| 31 | 3029 | 97.6 | 641 | 3 | US-09-897-778-339 | Sequence | 339, App |
| 32 | 3029 | 97.6 | 641 | 4 | US-10-007-700-339 | Sequence | 339, App |
| 33 | 3029 | 97.6 | 641 | 4 | US-10-117-982-339 | Sequence | 339, App |
| 34 | 3029 | 97.6 | 641 | 4 | US-10-313-986-339 | Sequence | 339, App |
| 35 | 3029 | 97.6 | 641 | 5 | US-10-775-972-339 | Sequence | 339, App |
| 36 | 3029 | 97.6 | 641 | 5 | US-10-922-124-339 | Sequence | 339, App |
| 37 | 3029 | 97.6 | 641 | 5 | US-10-623-155-339 | Sequence | 339, App |
| 38 | 3029 | 97.6 | 641 | 5 | US-10-469-469-282 | Sequence | 282, App |
| 39 | 3029 | 97.6 | 641 | 5 | US-10-469-469-298 | Sequence | 298, App |
| 40 | 3025 | 97.5 | 680 | 3 | US-09-735-705-342 | Sequence | 342, App |
| 41 | 3025 | 97.5 | 680 | 3 | US-09-850-716A-342 | Sequence | 342, App |
| 42 | 3025 | 97.5 | 680 | 3 | US-09-897-778-342 | Sequence | 342, App |
| 43 | 3025 | 97.5 | 680 | 4 | US-10-007-700-342 | Sequence | 342, App |
| 44 | 3025 | 97.5 | 680 | 4 | US-10-117-982-342 | Sequence | 342, App |
| 45 | 3025 | 97.5 | 680 | 4 | US-10-313-986-342 | Sequence | 342, App |
| 46 | 3025 | 97.5 | 680 | 5 | US-10-775-972-342 | Sequence | 342, App |
| 47 | 3025 | 97.5 | 680 | 5 | US-10-922-124-342 | Sequence | 342, App |
| 48 | 3025 | 97.5 | 680 | 5 | US-10-623-155-342 | Sequence | 342, App |
| 49 | 2995 | 96.5 | 680 | 4 | US-10-716-359-19 | Sequence | 19, Appl |
| 50 | 2995 | 96.5 | 680 | 6 | US-11-109-831-19 | Sequence | 19, Appl |
| 51 | 2421 | 78.0 | 461 | 3 | US-09-735-705-343 | Sequence | 343, App |
| 52 | 2421 | 78.0 | 461 | 3 | US-09-850-716A-343 | Sequence | 343, App |
| 53 | 2421 | 78.0 | 461 | 3 | US-09-897-778-343 | Sequence | 343, App |
| 54 | 2421 | 78.0 | 461 | 4 | US-10-007-700-343 | Sequence | 343, App |
| 55 | 2421 | 78.0 | 461 | 4 | US-10-117-982-343 | Sequence | 343, App |
| 56 | 2421 | 78.0 | 461 | 4 | US-10-313-986-343 | Sequence | 343, App |
| 57 | 2421 | 78.0 | 461 | 4 | US-10-716-359-17 | Sequence | 17, Appl |
| 58 | 2421 | 78.0 | 461 | 5 | US-10-775-972-343 | Sequence | 343, App |
| 59 | 2421 | 78.0 | 461 | 5 | US-10-922-124-343 | Sequence | 343, App |
| 60 | 2421 | 78.0 | 461 | 5 | US-10-623-155-343 | Sequence | 343, App |

OM protein - protein search, using sw model

December 27, 2006, 21:23:46; Search time 46 Seconds Run on:

(without alignments)

1246.430 Million cell updates/sec

US-10-716-359-16

Perfect score: 3104

1 MLYLENNAQTQFSEPQYTNL.....FNFDMDARRNKQQRIKEEGE 586 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

390055 segs, 97842647 residues Searched:

390055 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 500 summaries

Database : Published Applications AA New: *

1: /EMC Celerra SIDS3/ptodata/2/pubpaa/US09_NEW_PUB.pep:*

2: /EMC Celerra SIDS3/ptodata/2/pubpaa/US06_NEW_PUB.pep:*

3: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US07_NEW_PUB.pep:*

4: /EMC Celerra SIDS3/ptodata/2/pubpaa/US08 NEW_PUB.pep:*

5: /EMC Celerra SIDS3/ptodata/2/pubpaa/PCT_NEW_PUB.pep:*

6: /EMC Celerra SIDS3/ptodata/2/pubpaa/US10 NEW PUB.pep:*

/EMC Celerra SIDS3/ptodata/2/pubpaa/US11 NEW PUB.pep:*

8: /EMC Celerra SIDS3/ptodata/2/pubpaa/US60 NEW_PUB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | Query Match | Length | DB | ID | Description |
|---------------|--------|----------------|--------|----|-------------------|-------------------|
| 1 | 3033 | 97.7 | 641 | 7 | US-11-340-715-4 | Sequence 4, Appli |
| 2 | 2995 | 96.5 | 641 | 7 | US-11-340-715-11 | Sequence 11, Appl |
| 3 | 1813 | 58.4 | 448 | 7 | US-11-340-715-1 | Sequence 1, Appli |
| 4 | 1746 | 56.2 | 587 | 6 | US-10-489-730-2 | Sequence 2, Appli |
| 5 | 1741 | 56.1 | 636 | 7 | US-11-340-715-8 | Sequence 8, Appli |
| 6 | 1718.5 | 55.4 | 661 | 6 | US-10-489-730-11 | Sequence 11, Appl |
| 7 | 1420.5 | 45.8 | 450 | 6 | US-10-489-730-4 | Sequence 4, Appli |
| 8 | 1415.5 | 45.6 | 499 | 7 | US-11-340-715-6 | Sequence 6, Appli |
| 9 | 1256 | 40.5 | 426 | 6 | US-10-489-730-6 | Sequence 6, Appli |
| 10 | 1046 | 33.7 | 312 | 7 | US-11-340-715-7 | Sequence 7, Appli |
| 11 | 960.5 | 30.9 | 349 | 7 | US-11-340-715-9 | Sequence 9, Appli |
| 12 | 707.5 | 22.8 | 354 | 6 | US-10-538-066-763 | Sequence 763, App |
| 13 | 707.5 | 22.8 | 393 | 6 | US-10-538-066-367 | Sequence 367, App |
| 14 | 707.5 | 22.8 | 393 | 7 | US-11-315-777-9 | Sequence 9, Appli |
| 15 | 707.5 | 22.8 | 393 | 7 | US-11-340-715-3 | Sequence 3, Appli |
| 16 | 707.5 | 22.8 | 393 | 7 | US-11-009-357-6 | Sequence 6, Appli |
| 17 | 707.5 | 22.8 | 393 | 7 | US-11-319-873-9 | Sequence 9, Appli |

| 18 | 707.5 | 22.8 | 393 | 7 | UŞ-11-254-926-9 | Sequence 9, Appli |
|-----|-------|------|------|---|---------------------|-------------------|
| 19 | 707.5 | 22.8 | 393 | 7 | US-11-304-280-9 | Sequence 9, Appli |
| 20 | 707.5 | 22.8 | 393 | 7 | US-11-239-548-9 | Sequence 9, Appli |
| 21 | 707.5 | 22.8 | 425 | 7 | US-11-009-357-2 | Sequence 2, Appli |
| 22 | 702.5 | 22.6 | 393 | 6 | US-10-546-829-1 | Sequence 1, Appli |
| 23 | 702.5 | 22.6 | 393 | 7 | US-11-404-146-3 | Sequence 3, Appli |
| 24 | 699.5 | 22.5 | 393 | 7 | US-11-371-354-72585 | Sequence 72585, A |
| 25 | 316 | 10.2 | 102 | 6 | US-10-538-066-765 | Sequence 765, App |
| 26. | 291.5 | 9.4 | 100 | 6 | US-10-489-730-12 | Sequence 12, Appl |
| 27 | 285.5 | 9.2 | 100 | 6 | US-10-489-730-14 | Sequence 14, Appl |
| 28 | 284.5 | 9.2 | 100 | 6 | US-10-489-730-13 | Sequence 13, Appl |
| 29 | 133 | 4.3 | 2801 | 7 | US-11-375-359-120 | Sequence 120, App |
| 30 | 133 | 4.3 | 2896 | 7 | US-11-375-359-121 | Sequence 121, App |

OM nucleic - protein search, using frame plus n2p model Run on: December 27, 2006, 19:24:15; Search time 59 Seconds (without alignments) 4094.024 Million cell updates/sec US-10-716-359-4 Title: Perfect score: 3260 Sequence: 1 atgttgtacctggaaaacaa.....tcaaagaggagggggggtga 1761 BLOSUM62 Scoring table: Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0, Fgapext 7.0 6.0 , Delext 7.0 Delop 2589679 seqs, 457216429 residues Searched: Total number of hits satisfying chosen parameters: 5179358 Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ n2p.model -DEV=xlp -Q=/abss/ABSSWEB_spool/US10716359/runat_27122006_140406_325/app_query.fasta_1 -DB=A_Geneseq -QFMT=fastan -SUFFIX=n2p.rag -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=200000000 -HOST=abss02p -USER-US10716359_@CGN_1_1_440_@runat_27122006_140406_325 -NCPU=6 -ICPU=3 -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : A Geneseq 8:* 1: geneseqp1980s:* 2: geneseqp1990s:* 3: geneseqp2000s:* 4: geneseqp2001s:* 5: geneseqp2002s:*

6: geneseqp2003as:* 7: geneseqp2003bs:* 8: geneseqp2004s:* 9: geneseqp2005s:* 10: geneseqp2006s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

ક Result. Query No. Score Match Length DB ID Description

| | | | | | • | |
|----|--------|------|-----|-----|----------|----------------------|
| 1 | 3104 | 95.2 | 586 | 2 | AAY05956 | Aay05956 Human cel |
| 2 | 3104 | 95.2 | 586 | 5 | ABG95140 | Abg95140 Human onc |
| 3 | 3100 | 95.1 | 586 | 5 | ABG95132 | Abg95132 Human onc |
| 4 | 3087 | 94.7 | 586 | 3 | AAB11357 | Aab11357 Human p63 |
| 5 | 3087 | 94.7 | 586 | 5 | ABB74989 | Abb74989 Human p53 |
| 6 | 3087 | 94.7 | 586 | 5 | ABP61909 | Abp61909 Human lun |
| 7 | 3087 | 94.7 | 586 | 7 | ADH36992 | Adh36992 Human lun |
| 8 | 3087 | 94.7 | 586 | 8 | ADM56795 | Adm56795 Human lun |
| 9 | . 3087 | 94.7 | 586 | 8 | ADN04517 | Adn04517 Antipsori |
| 10 | 3087 | 94.7 | 586 | 9 | ADU98487 | Adu98487 Human p53 |
| 11 | 3087 | 94.7 | 586 | 9 | AEB10295 | Aeb10295 Cancer re |
| 12 | 3087 | 94.7 | 586 | 9 | AEE06149 | Aee06149 p53 tumor |
| 13 | 3080 | 94.5 | 586 | 2 | AAY41032 | · Aay41032 Human lun |
| 14 | 3080 | 94.5 | 586 | 3 | AAB11317 | Aab11317 Human lun |
| 15 | 3080 | 94.5 | 586 | 5 | ABB74949 | Abb74949 Human lun |
| 16 | 3080 | 94.5 | 586 | 5 | ABP61869 | Abp61869 Human lun |
| 17 | 3080 | 94.5 | 586 | 7 | ADE53447 | Ade53447 Human lun |
| 18 | 3080 | 94.5 | 586 | 7 | ADH36806 | Adh36806 Human lun |
| 19 | 3080 | 94.5 | 586 | 8 | ADM56609 | Adm56609 Human lun |
| 20 | 3080 | 94.5 | 586 | 8 | ADN89653 | Adn89653 Human lun |
| 21 | 3080 | 94.5 | 586 | . 9 | ADU98301 | Adu98301 Lung tumo |
| 22 | 3080 | 94.5 | 586 | 9 | AEB10109 | Aeb10109 Cancer re |
| 23 | 3080 | 94.5 | 586 | 9 | AEE05963 | Aee05963 Human lun |
| 24 | 3078 | 94.4 | 586 | 7 | ADA28428 | Ada28428 p53 tumou |
| 25 | 3077 | 94.4 | 586 | 7 | ADA28242 | Ada28242 Human lun |
| 26 | 3066 | 94.0 | 586 | 2 | AAY05962 | Aay05962 Mouse cel |
| 27 | 3033 | 93.0 | 641 | 2 | AAY45247 | Aay45247 Human p51 |
| 28 | 3033 | 93.0 | 641 | 2 | AAY05953 | Aay05953 Human cel |
| 29 | 3033 | 93.0 | 641 | 4 | AAB82129 | Aab82129 Human pro |
| 30 | 3033 | 93.0 | 641 | 5 | ABG95142 | · Abg95142 Human onc |
| 31 | 3033 | 93.0 | 687 | 7 | ADL22695 | Adl22695 Human dis |
| 32 | 3029 | 92.9 | 641 | 3 | AAB11358 | Aab11358 Human p63 |
| 33 | 3029 | 92.9 | 641 | 5 | ABG95136 | Abg95136 Human onc |
| 34 | 3029 | 92.9 | 641 | 5 | ABG95128 | Abg95128 Human onc |
| 35 | 3029 | 92.9 | 641 | 5 | ABB74990 | Abb74990 Human p53 |
| 36 | 3029 | 92.9 | 641 | 5 | ABP61910 | Abp61910 Human lun |
| 37 | 3029 | 92.9 | 641 | 7 | ADA28429 | Ada28429 p53 tumou |
| 38 | 3029 | 92.9 | 641 | 7 | ADH36993 | Adh36993 Human lun |
| 39 | 3029 | 92.9 | 641 | 8 | ADM56796 | Adm56796 Human lun |
| 40 | 3029 | 92.9 | 641 | 9 | ADU98488 | Adu98488 Human p53 |
| 41 | 3029 | 92.9 | 641 | 9 | AEB10296 | Aeb10296 Cancer re |
| 42 | 3029 | 92.9 | 641 | 9 | AEE06150 | Aee06150 p53 tumor |
| 43 | 3025 | 92.8 | 680 | 3 | AAY50997 | Aay50997 Human KET |
| 44 | 3025 | 92.8 | 680 | 3 | AAB11361 | Aab11361 Human p63 |
| 45 | 3025 | 92.8 | 680 | 5 | ABB74993 | Abb74993 Human p53 |

<!--EndFragment-->

OM nucleic - protein search, using frame_plus_n2p model

Run on: December 27, 2006, 19:26:20; Search time 107.5 Seconds

(without alignments)

4545.914 Million cell updates/sec

Title: US-10-716-359-4

Perfect score: 3260

Sequence: 1 atgttgtacctggaaaacaa.....tcaaagaggaggggggggtga 1761

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5
Ygapop 10.0 , Ygapext 0.5
Fgapop 6.0 , Fgapext 7.0
Delop 6.0 , Delext 7.0

Searched: 2849598 segs, 925015592 residues

Total number of hits satisfying chosen parameters: 5699196

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+_n2p.model -DEV=x1p

-Q=/abss/ABSSWEB_spool/US10716359/runat_27122006_140407_336/app_query.fasta_1

-DB=UniProt -QFMT=fastan -SUFFIX=n2p.rup -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -HOST=abss02p

-USER=US10716359 @CGN 1 1 891 @runat 27122006_140407_336 -NCPU=6 -ICPU=3

-NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120

-WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7

-YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database: UniProt 7.2:*

1: uniprot_sprot:*
2: uniprot trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| 8 | |
|------------------------------|-----------------------------|
| Result Query | • |
| No. Score Match Length DB ID | Description |
| | |
| 1 3033 93.0 680 1 P73L | _HUMAN Q9h3d4 homo sapien |
| 2 3026 92.8 582 2 Q3UV | I3_MOUSE Q3uvi3 mus musculu |
| 3 3017 92.5 582 2 Q5CZ | XO_MOUSE Q5czx0 mus musculu |
| 4 2995 91.9 680 1 P73L | MOUSE · 088898 mus musculu |
| 5 2991 91.7 680 1 P73L | RAT Q9jjp6 rattus norv |
| 6 2962 90.9 582 2 Q9DE | C7 CHICK Q9dec7 gallus gall |
| 7 2416 74.1 565 2 Q4S1 | 22 TETNG Q4s122 tetraodon n |
| 8 2344 71.9 576 2 Q8JF | E3_BRARE Q8jfe3 brachydanio |

| 9 | 2330 | 71.5 | 588 | 2 | Q8JHZ6 BRARE | Q8jhz6 | brachydanio |
|----|---------|------|-----|---|---------------|---------|-------------|
| 10 | 1877 | 57.6 | 393 | 2 | Q569E5 MOUSE | Q569e5 | mus musculu |
| 11 | 1826 | 56.0 | 457 | 2 | Q8JHZ5 BRARE | Q8jhz5 | brachydanio |
| 12 | 1777 | 54.5 | 641 | 2 | Q9W664_BARBU | Q9w664 | barbus barb |
| 13 | 1774 | 54.4 | 640 | 2 | Q801Z7 BRARE | Q801z7 | brachydanio |
| 14 | 1771 | 54.3 | 640 | 2 | Q6UNX2_BRARE | Q6unx2 | brachydanio |
| 15 | 1748.5 | 53.6 | 590 | 2 | Q9JJP1 MOUSE | Q9jjp1 | mus musculu |
| 16 | 1744.5 | 53.5 | 631 | 2 | Q9JJP2 MOUSE | Q9jjp2 | mus musculu |
| 17 | 1741 | 53.4 | 636 | 1 | P73_HUMAN | 015350 | homo sapien |
| 18 | 1730.5 | 53.1 | 365 | 2 | Q98SW0_XENLA | Q98sw0 | xenopus lae |
| 19 | 1720.5 | 52.8 | 637 | 1 | P73_CERAE | ·Q9xsk8 | cercopithec |
| 20 | 1649.5 | 50.6 | 434 | 2 | Q6DG24_BRARE | Q6dg24 | brachydanio |
| 21 | 1562 | 47.9 | 514 | 2 | Q9CU77_MOUSE | Q9cu77 | mus musculu |
| 22 | 1,507.5 | 46.2 | 494 | 2 | Q3UT91_MOUSE | Q3ut91 | mus musculu |
| 23 | 1481 | 45.4 | 662 | 2 | Q4S837_TETNG | Q4s837 | tetraodon n |
| 24 | 1464 | 44.9 | 555 | 2 | Q5KQU6_BRARE | Q5kqu6 | brachydanio |
| 25 | 1459.5 | 44.8 | 497 | 2 | Q9WUJ0_MOUSE | Q9wuj0 | mus musculu |
| 26 | 950.5 | 29.2 | 621 | 2 | Q9NGC7_MYAAR | Q9ngc7 | mya arenari |
| 27 | 912 | 28.0 | 591 | 2 | Q8T7V3_SPISO` | Q8t7v3 | spisula sol |
| 28 | 912 | 28.0 | 596 | 2 | Q6WG19_SPISO | | spisula sol |
| 29 | 912 | 28.0 | 596 | 2 | Q6WG20_SPISO | Q6wg20 | spisula sol |
| 30 | 898.5 | 27.6 | 564 | 2 | Q27937 LOLFO | Q27937 | loligo forb |

<!--StartFragment-->

GenCore version 5.1.9

Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM nucleic - protein search, using frame plus n2p model

Run on: December 27, 2006, 19:36:35; Search time 13.3 Seconds

(without alignments)

3821.903 Million cell updates/sec

Title: US-10-716-359-4

Perfect score: 3260

Sequence: 1 atgttgtacctggaaaacaa.....tcaaagaggagggggggtga 1761

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5
Ygapop 10.0 , Ygapext 0.5
Fgapop 6.0 , Fgapext 7.0
Delop 6.0 , Delext 7.0

Searched: 283416 segs, 96216763 residues

Total number of hits satisfying chosen parameters: 566832

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+_n2p.model -DEV=xlp

-Q=/abss/ABSSWEB spool/US10716359/runat 27122006 140410 379/app_query.fasta_1

-DB=PIR -QFMT=fastan -SUFFIX=n2p.rpr -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=200000000 -HOST=abss02p

-USER=US10716359_@CGN_1_1_92_@runat_27122006_140410_379 -NCPU=6 -ICPU=3

-NO_MMAP -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120

-WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7

-YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database: PIR_80:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result | | % Ouerv | | | | | | | | |
|--------|-------|------------|--------|----|--------|---|-----------|-----------|-------|------|
| No. | Score | 1 | Length | DB | ID | | | Descripti | ion | |
| 1 | 1774 | 54.4 | 640 | 2 | JC7989 | | · | p73 prote | ein - | zebr |
| 2 | 839.5 | 25.8 | 396 | 1 | JH0631 | | • | cellular | tumor | ant |
| 3 | 773.5 | 23.7 | 367 | 1 | S02193 | | | cellular | tumor | ant |
| 4 | 762 | . 23.4 | 363 | 1 | A29376 | | | cellular | tumor | ant |
| 5 | 747.5 | 22.9 | 386 | 1 | S51648 | | | cellular | tumor | ant |
| 6 | 736.5 | 22.6 | 391 | 1 | S02192 | • | • | cellular | tumor | ant |
| | | | | | | | | | | |

| 7 | 726 | 22.3 | 393 | 1 | DNHU53 | |
|----|-------|------|-------|---|--------|--|
| 8 | 719.5 | 22.1 | 393 | 1 | S06594 | |
| 9 | 712 | 21.8 | 396 | 1 | JH0633 | |
| 10 | 709.5 | 21.8 | 390 | 1 | DNMS53 | |
| 11 | 701.5 | 21.5 | 391 | 2 | JC6193 | |
| 12 | 697.5 | 21.4 | 393 | 2 | JC6176 | |
| 13 | 692.5 | 21.2 | 381 | 2 | S38824 | |
| 14 | 259.5 | 8.0 | 77 | 2 | I46226 | |
| 15 | 161 | 4.9 | 925 | 2 | T19361 | |
| 16 | 152.5 | 4.7 | 3570 | 2 | T45025 | |
| 17 | 151.5 | 4.6 | 3020 | 2 | A43932 | |
| 18 | 151 | 4.6 | 2715 | 2 | T13049 | |
| 19 | 150 | 4.6 | 1006 | 2 | T42731 | |
| 20 | 150 | 4.6 | 1188 | 2 | S49915 | |
| 21 | 147.5 | 4.5 | 620 | 2 | S06733 | |
| 22 | 144.5 | 4.4 | 862 | 2 | B53689 | |
| 23 | 142.5 | 4.4 | 2359 | 2 | T03094 | |
| 24 | 140.5 | 4.3 | 1199 | 2 | A40670 | |
| 25 | 140.5 | 4.3 | 2232 | 2 | T34434 | |
| 26 | 140 | 4.3 | 847 | 2 | F96531 | |
| 27 | 139 . | 4.3 | 1285 | 2 | T14171 | |
| 28 | 139 | 4.3 | 1613 | 2 | S39059 | |
| 29 | 138 | 4.2 | 1510 | 2 | T33100 | |
| 30 | 136 | 4.2 | 400 | 2 | S58222 | |
| 31 | 136 | 4.2 | 650 | 2 | T04487 | |
| 32 | 135.5 | 4.2 | 891 | 2 | G84693 | |
| 33 | 135.5 | 4.2 | 2649 | 2 | T51023 | |
| 34 | 135 | 4.1 | 975 | 2 | S33121 | |
| 35 | 135 | 4.1 | .1240 | 2 | JC5209 | |
| 36 | 135 | 4.1 | 1242 | 2 | JS0670 | |
| 37 | 134.5 | 4.1 | 628 | 2 | JQ0110 | |
| 38 | 134.5 | 4.1 | 901 | 2 | JC6093 | |
| 39 | 134 | 4.1 | 1520 | 1 | TVFFA | |
| 40 | 134 | 4.1 | 1791 | 2 | T24089 | |
| 41 | 133.5 | 4.1 | 477 | 2 | Т46304 | |
| 42 | 133.5 | 4.1 | 964 | 2 | T41547 | |
| 43 | 133 | 4.1 | 817 | 2 | S51342 | |
| 44 | 133 | 4.1 | 1054 | 2 | T43226 | |
| | | | | _ | | |

1081 2 S66736

cellular tumor ant cellular tumor ant cellular tumor ant cellular tumor ant tumor suppressor p tumor suppressor p cellular tumor ant cellular tumor ant hypothetical prote mucin MUC5B, trach mucin 2 precursor, eyelid - fruit fly atrophin-1 related extensin-like prot hydroxyproline-ric homeotic protein C A-kinase anchor pr nuclear envelope p hypothetical prote hypothetical prote ataxin-2 - mouse protein BRG1 - hum hypothetical prote PQ-rich protein hypothetical prote probable proline-r hypothetical prote homeotic protein C insulin receptor s insulin receptor s hypothetical 69K p dead ringer nuclea protein-tyrosine k hypothetical prote hypothetical prote hypothetical prote verprolin - yeast translation initia transcription acti

<!--EndFragment-->

133

4.1

45

OM nucleic - protein search, using frame plus n2p model

December 27, 2006, 19:54:31; Search time 12.8 Seconds Run on:

(without alignments)

3612.690 Million cell updates/sec

US-10-716-359-4 Title:

Perfect score: 3260

1 atgttqtacctqqaaaacaa.....tcaaagaggagggggggtga 1761 Sequence:

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 . Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0, Fgapext 7.0 6.0 , Delext 7.0 Delop

650591 seqs, 87530628 residues Searched:

Total number of hits satisfying chosen parameters: 1301182

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

- -MODEL=frame+ n2p.model -DEV=xlp
- -Q=/abss/ABSSWEB spool/US10716359/runat 27122006 140412 434/app_query.fasta_1
- -DB=Issued Patents AA -QFMT=fastan -SUFFIX=n2p.rai -MINMATCH=0.1 -LOOPCL=0
- -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi
- -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15
- -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -HOST=abss02p -USER=US10716359_@CGN_1_1_84_@runat_27122006_140412_434 -NCPU=6
- -ICPU=3 -NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120
- -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7
- -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Issued_Patents_AA:* Database :

- 1: /EMC Celerra SIDS3/ptodata/2/iaa/5_COMB.pep:*
- 2: /EMC_Celerra_SIDS3/ptodata/2/iaa/6_COMB.pep:*
- 3: /EMC Celerra SIDS3/ptodata/2/iaa/7 COMB.pep:*
- 4: /EMC_Celerra_SIDS3/ptodata/2/iaa/H_COMB.pep:*
- 5: /EMC_Celerra_SIDS3/ptodata/2/iaa/PCTUS_COMB.pep:*
- 6: /EMC_Celerra_SIDS3/ptodata/2/iaa/RE_COMB.pep:*
- /EMC Celerra SIDS3/ptodata/2/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | Query Match | Length | DB | ID | Description |
|---------------|-------|----------------|--------|----|-------------------|-------------------|
| 1 | 3104 | 95.2 | 586 | 2 | US-09-538-106-16 | Sequence 16, Appl |
| 2 | 3104 | 95.2 | 586 | 3 | US-09-529-583-16 | Sequence 16, Appl |
| 3 | 3087 | 94.7 | 586 | 2 | US-09-643-597-338 | Sequence 338, App |

| 4 | 3087 | 94.7 · | 586 | 2 | US-09-542-615A-338 | Sequence | 338, App |
|------|------|--------|-----|---|--------------------|--------------|----------|
| 5 | 3087 | 94.7 | 586 | 2 | US-09-606-421B-338 | | 338, App |
| 6 | 3087 | 94.7 | 586 | 2 | US-09-630-940B-338 | Sequence | 338, App |
| 7 | 3087 | 94.7 | 586 | 2 | US-10-007-700-338 | Sequence | 338, App |
| 8 | 3080 | 94.5 | 586 | 2 | US-09-643-597-152 | Sequence | 152, App |
| 9 | 3080 | 94.5 | 586 | 2 | US-09-480-884A-152 | Sequence | 152, App |
| 10 | 3080 | 94.5 | 586 | 2 | US-09-542-615A-152 | Sequence | 152, App |
| 11 | 3080 | 94.5 | 586 | 2 | US-09-606-421B-152 | Sequence | 152, App |
| 12 | 3080 | 94.5 | 586 | 2 | US-09-221-107-152 | Sequence | 152, App |
| 13 | 3080 | 94.5 | 586 | 2 | US-09-466-396A-152 | Sequence | 152, App |
| · 14 | 3080 | 94.5 | 586 | 2 | US-09-476-496A-152 | Sequence | 152, App |
| 15 | 3080 | 94.5 | 586 | 2 | US-09-630-940B-152 | Sequence | 152, App |
| 16 | 3080 | 94.5 | 586 | 2 | US-09-285-479-152 | Sequence | 152, App |
| 17 | 3080 | 94.5 | 586 | 2 | US-10-007-700-152 | Sequence | 152, App |
| 18 | 3066 | 94.0 | 586 | 2 | US-09-538-106-22 | Sequence | 22, Appl |
| 19 | 3066 | 94.0 | 586 | 3 | US-09-529-583-22 | Sequence | 22, Appl |
| 20 | 3033 | 93.0 | 641 | 2 | US-09-538-106-13 | Sequence | 13, Appl |
| 21 | 3033 | 93.0 | 641 | 3 | US-09-529-583-13 | Sequence | 13, Appl |
| 22 | 3029 | 92.9 | 641 | 2 | US-09-643-597-339 | Sequence | 339, App |
| 23 | 3029 | 92.9 | 641 | 2 | US-09-542-615A-339 | Sequence | 339, App |
| 24 | 3029 | 92.9 | 641 | 2 | US-09-606-421B-339 | Sequence | 339, App |
| 25 | 3029 | 92.9 | 641 | 2 | US-09-630-940B-339 | Sequence | 339, App |
| 26 | 3029 | 92.9 | 641 | 2 | US-10-007-700-339 | Sequence | 339, App |
| 27 | 3025 | 92.8 | 680 | 2 | US-09-643-597-342 | Sequence | 342, App |
| 28 | 3025 | 92.8 | 680 | 2 | US-09-542-615A-342 | Sequence | 342, App |
| 29 | 3025 | 92.8 | 680 | 2 | US-09-606-421B-342 | | 342, App |
| 30 | 3025 | 92.8 | 680 | 2 | US-09-630-940B-342 | Sequence | 342, App |
| 31 | 3025 | 92.8 | 680 | 2 | US-10-007-700-342 | Sequence | 342, App |
| 32 | 2995 | 91.9 | 680 | 2 | US-09-538-106-19 | Sequence | 19, Appl |
| 33 | 2995 | 91.9 | 680 | 3 | US-09-529-583-19 | | 19, Appl |
| 34 | 2421 | 74.3 | 461 | 2 | US-09-643-597-343 | Sequence | 343, App |
| 35 | 2421 | 74.3 | 461 | 2 | US-09-542-615A-343 | Sequence | 343, App |
| 36 | 2421 | 74.3 | 461 | 2 | US-09-606-421B-343 | Sequence | 343, App |
| · 37 | 2421 | 74.3 | 461 | 2 | US-09-630-940B-343 | Sequence | 343, App |
| 38 | 2421 | 74.3 | 461 | 2 | US-09-538-106-17 | | 17, Appl |
| 39 | 2421 | 74.3 | 461 | 2 | US-10-007-700-343 | Sequence | 343, App |
| 40 | 2421 | 74.3 | 461 | 3 | US-09-529-583-17 | | 17, Appl |
| 41 | 2398 | 73.6 | 461 | 2 | US-09-538-106-23 | | 23, Appl |
| 42 | 2398 | 73.6 | 461 | 3 | US-09-529-583-23 | | 23, Appl |
| 43 | 2350 | 72.1 | 516 | 2 | US-09-643-597-344 | | 344, App |
| 44 | 2350 | 72.1 | 516 | 2 | US-09-542-615A-344 | - | 344, App |
| 45 | 2350 | 72.1 | 516 | 2 | US-09-606-421B-344 | Sequence | 344, App |

OM nucleic - protein search, using frame plus_n2p model

Run on: December 27, 2006, 20:28:36; Search time 58.9 Seconds

(without alignments)

4154.778 Million cell updates/sec

Title: US-10-716-359-4

Perfect score: 3260

Sequence: 1 atgttgtacctggaaaacaa.....tcaaagaggaggggggggtga 1761

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5
Ygapop 10.0 , Ygapext 0.5
Fgapop 6.0 , Fgapext 7.0
Delop 6.0 , Delext 7.0

Searched: 2097797 segs, 463214858 residues

Total number of hits satisfying chosen parameters: 4195594

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+_n2p.model -DEV=xlp

-Q=/abss/ABSSWEB spool/US10716359/runat 27122006 140422 629/app_query.fasta_1

-DB=Published_Applications_AA_Main -QFMT=fastan -SUFEIX=n2p.rapbm

-MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1

-MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct

-THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext

-HEAPSIZE=500 -MINLEN=0 -MAXLEN=200000000 -HOST=abss02p

-USER=US10716359_@CGN_1_1_446_@runat_27122006_140422_629 -NCPU=6 -ICPU=3

-NO MMAP -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120

-WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7

-YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database: Published Applications AA Main:*

1: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US07_PUBCOMB.pep:*

2: /EMC_Celerra_SIDs3/ptodata/2/pubpaa/US08_PUBCOMB.pep:*

3: /EMC Celerra SIDS3/ptodata/2/pubpaa/US09 PUBCOMB.pep:*

4: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US10A PUBCOMB.pep:*

5: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US10B_PUBCOMB.pep:*

6: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US11_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | % Query Match | Length | DB | ID | Description |
|---------------|--------------|---------------------|------------|----|---------------------------------------|--|
| 1 | 3104 | 95.2 | | _ | US-10-716-359-16 | Sequence 16, Appl |
| 2 3 | 3104 3104 | 95.2 95.2 | 586 586 | - | US-10-469-469-306 US-11-109-831-16 | Sequence 306, App Sequence 16, Appl |

| 4 | 3100 | 95.1 | 586 | 5 | US-10-469-469-290 | Sequence | 290, App |
|----|------|------|-----|---|----------------------------|------------|----------|
| 5 | 3087 | 94.7 | 586 | 3 | US-09-735-705-338 | Sequence | 338, App |
| 6 | 3087 | 94.7 | 586 | 3 | US-09-850-716A-338 | Sequence | 338, App |
| 7 | 3087 | 94.7 | 586 | 3 | US-09-897-778-338 | Sequence | 338, App |
| 8 | 3087 | 94.7 | 586 | 4 | US-10-007-700-338 | Sequence | 338, App |
| 9 | 3087 | 94.7 | 586 | 4 | US-10-117-982-338 | Sequence | 338, App |
| 10 | 3087 | 94.7 | 586 | 4 | US-10-313-986-338 | Sequence | 338, App |
| 11 | 3087 | 94.7 | 586 | 5 | US-10-775-972-338 | Sequence | 338, App |
| 12 | 3087 | 94.7 | 586 | 5 | US-10-922-124-338 | Sequence | 338, App |
| 13 | 3087 | 94.7 | 586 | 5 | US-10-623-155-338 | Sequence | 338, App |
| 14 | 3080 | 94.5 | 586 | 3 | US-09-735-705-152 | Sequence | 152, App |
| 15 | 3080 | 94.5 | 586 | 3 | US-09-850-716A-152 | Sequence | 152, App |
| 16 | 3080 | 94.5 | 586 | 3 | US-09-897-778-152 | | 152, App |
| 17 | 3080 | 94.5 | 586 | 3 | US-09-466-396A-152 | Sequence | 152, App |
| 18 | 3080 | 94.5 | 586 | 4 | US-10-007-700-152 | | 152, App |
| 19 | 3080 | 94.5 | 586 | 4 | US-10-117-982-152 | , Sequence | 152, App |
| 20 | 3080 | 94.5 | 586 | 4 | US-10-313-986-152 | Sequence | 152, App |
| 21 | 3080 | 94.5 | 586 | 5 | US-10-775-972-152 | Sequence | 152, App |
| 22 | 3080 | 94.5 | 586 | 5 | US-10-922-124-152 | Sequence | 152, App |
| 23 | 3080 | 94.5 | 586 | 5 | US-10-623-155-152 | Sequence | 152, App |
| 24 | 3066 | 94.0 | 586 | 4 | US-10-716-359-22 | Sequence | 22, Appl |
| 25 | 3066 | 94.0 | 586 | 6 | US-11-109-831-22 | Sequence | 22, Appl |
| 26 | 3033 | 93.0 | 641 | 4 | US-10-716-359-13 | Sequence | 13, Appl |
| 27 | 3033 | 93.0 | 641 | 5 | US-10-469-469-310 | Sequence | 310, App |
| 28 | 3033 | 93.0 | 641 | 6 | US-11-109-831-13 | Sequence | 13, Appl |
| 29 | 3029 | 92.9 | 641 | 3 | US-09-735-705-339 | Sequence | 339, App |
| 30 | 3029 | 92.9 | 641 | 3 | US-09-850-716A-339 | Sequence | 339, App |
| 31 | 3029 | 92.9 | 641 | 3 | US-09-897-778-339 | Sequence | 339, App |
| 32 | 3029 | 92.9 | 641 | 4 | US-10-007-700-339 | Sequence | 339, App |
| 33 | 3029 | 92.9 | 641 | 4 | US-10-117-982-339 | Sequence | 339, App |
| 34 | 3029 | 92.9 | 641 | 4 | US-10-313-986-339 | Sequence | 339, App |
| 35 | 3029 | 92.9 | 641 | 5 | US-10-775-972-339 | Sequence | 339, App |
| 36 | 3029 | 92.9 | 641 | 5 | US-10-922-124-339 | Sequence | 339, App |
| 37 | 3029 | 92.9 | 641 | 5 | US-10-623-155-339 | Sequence | 339, App |
| 38 | 3029 | 92.9 | 641 | 5 | US-10-469-469-282 | Sequence | 282, App |
| 39 | 3029 | 92.9 | 641 | 5 | US-10-469-469 - 298 | Sequence | 298, App |
| 40 | 3025 | 92.8 | 680 | 3 | US-09-735-705-342 | Sequence | 342, App |
| 41 | 3025 | 92.8 | 680 | 3 | US-09-850-716A-342 | Sequence | 342, App |
| 42 | 3025 | 92.8 | 680 | 3 | US-09-897-778-342 | Sequence | 342, App |
| 43 | 3025 | 92.8 | 680 | 4 | US-10-007-700-342 | Sequence | 342, App |
| 44 | 3025 | 92.8 | 680 | 4 | US-10-117-982-342 | Sequence | 342, App |
| 45 | 3025 | 92.8 | 680 | 4 | US-10-313-986-342 | Sequence | 342, App |

OM nucleic - protein search, using frame_plus_n2p model

December 27, 2006, 20:35:01; Search time 13.7 Seconds Run on:

(without alignments)

3773.012 Million cell updates/sec

US-10-716-359-4 Title:

Perfect score: 3260

Sequence:

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0; Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 6.0 , Delext 7.0 Delop

390055 segs, 97842647 residues Searched:

Total number of hits satisfying chosen parameters: 780110

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+ n2p.model: -DEV=xlp

-Q=/abss/ABSSWEB spool/US10716359/runat 27122006 140426 732/app query.fasta_1

-DB=Published_Applications_AA_New -QFMT=fastan -SUFFIX=n2p.rapbn -MINMATCH=0.1

-LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62

-TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100

-THR MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0

-MAXLEN=2000000000 -HOST=abss02p

-USER=US10716359 @CGN 1 1 94 @runat 27122006 140426 732 -NCPU=6 -ICPU=3

-NO MMAP -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120

-WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7

-YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : Published Applications AA New: *

1: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US09_NEW_PUB.pep:*

/EMC_Celerra_SIDS3/ptodata/2/pubpaa/US06_NEW_PUB.pep:* 3: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US07_NEW_PUB.pep:*

4: /EMC Celerra SIDS3/ptodata/2/pubpaa/US08_NEW_PUB.pep:*

5: /EMC Celerra SIDS3/ptodata/2/pubpaa/PCT NEW PUB.pep:*

6: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US10_NEW_PUB.pep:*

7: /EMC Celerra SIDS3/ptodata/2/pubpaa/US11 NEW PUB.pep:*

8: /EMC Celerra SIDS3/ptodata/2/pubpaa/US60_NEW_PUB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| Result No. | Score | Query Match | Length | DB | ID | Description |
|---------------|-------|----------------|--------|--------------|-----------------|-------------------|
| 1 | 3033 | 93.0 | 641 | 7 | US-11-340-715-4 | Sequence 4, Appli |

| 2 | 2995 | 91.9 | 641 | 7 | US-11-340-715-11 | Sequence 11, Appl |
|----------|--------|------|------|-----|----------------------|-------------------|
| 3 | 1813 | 55.6 | 448 | 7 | US-11-340-715-1 | Sequence 1, Appli |
| 4 | 1746 | 53.6 | 587 | 6 | US-10-489-730-2 | Sequence 2, Appli |
| 5 | 1741 | 53.4 | 636 | 7 | US-11-340-715-8 | Sequence 8, Appli |
| 6 | 1718.5 | 52.7 | 661 | 6 | US-10-489-730-11 | Sequence 11, Appl |
| 7 | 1420.5 | 43.6 | 450 | 6 | US-10-489-730-4 | Sequence 4, Appli |
| 8 | 1420.5 | 43.4 | 499 | 7 | US-11-340-715-6 | Sequence 6, Appli |
| 9 | 1293.5 | 39.7 | 426 | 6 | US-10-489-730-6 | Sequence 6, Appli |
| 10 | 1046 | 32.1 | 312 | 7 | US-11-340-715-7 | Sequence 7, Appli |
| | 960.5 | 29.5 | 349 | 7 | US-11-340-715-7 | Sequence 9, Appli |
| 11 12 | | | 393 | 7 | US-11-340-715-3 | Sequence 3, Appli |
| | 726 | 22.3 | | 7 | | |
| 13. | | 22.3 | 393 | | US-11-009-357-6 | Sequence 6, Appli |
| 14 | 726 | 22.3 | 425 | 7 | US-11-009-357-2 | Sequence 2, Appli |
| 15 | 721 | 22.1 | 393 | 6 | US-10-546-829-1 | Sequence 1, Appli |
| 16 | 719 | 22.1 | 393 | 6 | US-10-538-066-367 | Sequence 367, App |
| 17 | 719 | 22.1 | 393 | 7 | US-11-315-777-9 | Sequence 9, Appli |
| 18 | 719 | 22.1 | 393 | 7 | US-11-319-873-9 | Sequence 9, Appli |
| 19 | 719 | 22.1 | 393 | 7 | US-11-254-926-9 | Sequence 9, Appli |
| 20 | 719 | 22.1 | 393 | 7 | US-11-304-280-9 | Sequence 9, Appli |
| 21 | 719 | 22.1 | 393 | 7 | US-11-239-548-9 | Sequence 9, Appli |
| 22 | 718 | 22.0 | 393 | 7 - | = | Sequence 72585, A |
| 23 | 714 | 21.9 | 393 | 7 | US-11-404-146-3 | Sequence 3, Appli |
| 24 | 713 | 21.9 | 354 | 6 | US-10-538-066-763 | Sequence 763, App |
| 25 | 316 | 9.7 | 102 | 6 | US-10-538-066-765 | Sequence 765, App |
| 26 | 291.5 | 8.9 | 100 | 6 | US-10-489-730-12 | Sequence 12, Appl |
| 27 | 285.5 | 8.8 | 100 | 6 | US-10-489-730-14 | Sequence 14, Appl |
| 28 | 284.5 | 8.7 | 100 | 6 | US-10-489-730-13 | Sequence 13, Appl |
| 29 | 163 | 5.0 | 2240 | 6 | US-10-544-731-4 | Sequence 4, Appli |
| 30 | 163 | 5.0 | 2258 | 6 | US-10-544-731-3 | Sequence 3, Appli |
| 31 | 163 | 5.0 | 2264 | 6 | US-10-544-731-5 | Sequence 5, Appli |
| 32 | 154.5 | 4.7 | 634 | 7 | US-11-293-697-3340 | Sequence 3340, Ap |
| 33 | 149.5 | 4.6 | 693 | 7 | US-11-293-697-3849 | Sequence 3849, Ap |
| 34 | 143.5 | 4.4 | 1113 | 7 | US-11-056-355B-77057 | Sequence 77057, A |
| 35 | 142.5 | 4.4 | 1690 | 6 | US-10-516-780-28 | Sequence 28, Appl |
| 36 | 141 | 4.3 | 690 | 6 | US-10-449-902-41507 | Sequence 41507, A |
| 37 | 140.5 | 4.3 | 312 | 6 | US-10-953-349-31493 | Sequence 31493, A |
| 38 | 140.5 | 4.3 | 312 | 7 | US-11-056-355B-68325 | Sequence 68325, A |
| 39 | 140 | 4.3 | 691 | 7 | US-11-056-355B-79590 | Sequence 79590, A |
| 40 | 140 | 4.3 | 718 | 7 | US-11-056-355B-79589 | Sequence 79589, A |
| 41 | 140 | 4.3 | 847 | 7 | US-11-056-355B-79588 | Sequence 79588, A |
| 42 | 138.5 | 4.2 | 5178 | 6 | US-10-700-439-178 | Sequence 178, App |
| 43 | 138.5 | 4.2 | 5179 | 6 | US-10-541-749-151 | Sequence 151, App |
| 44 | 138.5 | 4.2 | 5179 | 7 | US-11-105-233-185 | Sequence 185, App |
| 45 | 138 | 4.2 | 934 | 6 | US-10-449-902-41077 | Sequence 41077, A |
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